

# Erythrocyte Catalase Activities in Alcohol Consumption, Medications and Some Diseases

Ismail Temel\*, Elif Özerol\*, Aysun Bay\*, Ahmet Çigli\*, Ömer Akyol\*,

\*Inönü Üniversitesi, Tip Fakültesi, Biyokimya AD, Malatya

Catalase (CAT) has a catalytic role in the decomposition of hydrogen peroxide  $(H_2O_2)$  and a peroxidic role in which the peroxide is utilized to oxidize a range of H donors. The objective of the present study was to determine whether alcohol consumption, medications, and diseases have an effect in the erythrocyte on CAT activity or not. For this purpose, the subjects were divided into three different groups with their unique criteria: 1- alcoholic and nonalcoholic, 2- subjects under medication and no medication, and 3- subjects with leukemia, hepatitis, diabetes mellitus, and heart diseases.

CAT activity was measured by the method of Aebi. The hemoglobin levels were determined by Olympus AU-600 auto analyzer. Hematological parameters such as MCH, HCT, MCHC, MCV, and RBC were studied by Coulter STKS instrument. There was no effect of medication on CAT activity  $3155\pm1039$  k/g Hb for subjects under medication and  $3051\pm956$  for other subjects (p>0.05). When CAT activities of the disease groups with leukemic, cardiac, hepatitis, and diabetic were compared to the control group, there were no significant differences between them. We found that enzyme activity was affected significantly by only alcohol consumption. CAT activities were  $3059\pm958$  k/g Hb in nonalcoholic subjects and  $3644\pm984$  k/g Hb in alcoholic subjects (p<0.03).

Key words: Catalase - Red Blood Cell - Leukemia - Cardiac Disease - Hepatitis - Diabetes Mellitus

### Alkol Tüketimi, Ilaç Kullanimi ve Bazi Hastaliklarda Eritrosit Katalaz Aktiviteleri

Katalaz (CAT) hidrojen peroksidin parçalanmasında katalitik rol oynarken, peroksidin hidrojen donörüne okside edilmesinde peroksidik bir rol oynar. Bu çalismada alkol tüketimi, ilaçlar ve hastaliklarin eritrosit katalaz aktivitesi üzerine etkisini arastirmak amaçlanmistir. Bu amaç için denekler 3 farkli gruba ayrıldılar; 1- alkol alan ve almayan 2- ilaç alan ve almayan 3- lösemi, hepatit, diabetes mellitus ve kardiyak hastaliklari olanlar. Katalaz aktivitesi Aebi metodu ile ölçüldü. Hemoglobin düzeyleri Olympus AU-600 otoanalizörüyle tayin edildi.

Katalaz aktivitesi Aebi metodu ile olçuldu. Hemoglobin dužeyleri Olympus AU-600 otoanalizoruyle tayın edildi. MCH, HCT, MCHC, MCV ve RBC gibi hematolojik degerler Coulter STKS cihaziyla çalisildi. Ilaç alan ve almayanlarin katalaz aktivitesinde bir farklilik gözlenmedi (3155 ? 1039 k/g Hb ve 3051 ? 956 k/g Hb, p>.0,05). Saglikli kisilerdeki ile lösemililerin, kalp hastalarinin, hepatitlilerin ve diabetes mellituslularin katalaz aktiviteleri arasında fark bulunamadi. Alkol ile enzim aktivitesi belirgin bir sekilde artti. CAT aktiviteleri alkol almayanlarda 3059 ? 958 k/g Hb ve alkol alanlarda 3644 ? 984 k/g Hb olarak bulundu (p<0,03).

Anahtar kelimeler: Katalaz, Eritrosit, Lösemi, Kalp Hastaligi, Hepatit, Diyabet

## **INTRODUCTION**

Free oxygen radicals produced by normal aerobic metabolism have been implicated in several pathophysiological mammalian processes: Mammalian erythrocytes have large amounts of CAT, which is a heme-containing enzyme that catalyses the conversion of hydrogen peroxide to water and oxygen. <sup>1-3</sup> Catalase (CAT E.C.1.11.1.6) has a dual functional role; a true catalytic role in the decomposition of hydrogen peroxide ( $H_2O_2$ ) and a peroxidic role in which the peroxide is utilized to oxidize a range of H donors. It is widely distributed in the body compartments, tissues, and cells. Erythrocytes appear to have high CAT activity compared to other cells because of greatly exposure of erythrocytes to molecular oxygen. Hepatic CAT activities also increase in experimental endotoxemia and hepatitis.<sup>4,5</sup> In alcoholism, hepatic catalase activities decline, though as aforementioned, it appears to be altered in iron-overload.<sup>6,7</sup> A wide range of studies support the protective effect of dietary antioxidants for reducing the risk of

cardiovascular disease.<sup>8</sup> Reactive oxygen species (ROS) appear to be involved in both the development and later complications of diabetes. Moreover, evidence from both animal studies and humans suggest that antioxidant defenses become compromised prior to the development of diabetes.<sup>9-13</sup>

Red cell antioxidant enzymes have been recently studied in malignant lymphomas and results are controversial.<sup>14</sup> The components of the blood antioxidant systems (superoxide dismutase, catalase, ceruloplasmin, glutathione system) take a direct part in molecular mechanisms of the body adaptation under conditions of viral hepatitis infections.<sup>15</sup>

Changes in the antioxidant system of red blood cells may be recorded in chronic liver diseases (persistent and active hepatitis, liver cirrhosis). The findings in the literature were the activation of SOD and glutathione reductase; reduction of the activity of total and membrane-bound catalase the content of reduced glutathione.<sup>16</sup> On the other hand, reperfusion of ischemic heart causes the generation of free radicals, and these radicals play an important role in post-ischemic tissue damage. These free radicals are removed by scavenger enzymes and antioxidants in the cell.<sup>17</sup>

In a study, it was found that the oxygen free radical reaction in alcohol abusers was pathologically exacerbated and the balance between oxidation and antioxidation was seriously disturbed.<sup>18</sup> Since free radicals and peroxides seem to be involved in the toxicity of alcoholics. Several authors have examined the variations of blood activities of antioxidant enzymes in alcoholics, but published results are somewhat conflicting. Variations of blood antioxidant enzymes observed in patients were of limited amplitude and do not allow the use of either of them as markers of alcohol abuse.<sup>19</sup> Guemouri et al. observed the strong effects in therapy by antidepressants or thyroid hormones. Intake of some drugs (e.g., anti-inflammatory agents, antidepressants, and thyroid hormones) modifies activity of some of the three enzymes (SOD, GPX, and CAT).<sup>20</sup> The behavior of these metabolic parameters reveals the complexity of the diabetic red blood cell metabolism and in addition underlines the fact that the diabetic erythrocytes being less protected from the oxidant agents, has a reduced mean survival as has been evidentiated by some authors.<sup>21</sup>

GPX, reduced glutathione (GSH), SOD, and CAT were measured in homogeneous group of patients with untreated hairy cell leukemia and normal controls. GPX, CAT, and SOD activities were significantly lower in patients than in normals. Taken together, these data suggest a decreased activity of red cell antioxidant enzymes in hairy cell leukemia and support a pluripotent stem cell defect of these abnormalities.<sup>14</sup> Therefore, the objective of the present study was to determine whether alcohol consumption, medications, and diseases affect the erythrocyte CAT activity or not.

## MATERIALS AND METHODS

We prospectively selected patient admitted to Turgut Ozal Medical Center with a clinical diagnosis of leukemia, hepatitis, diabetes mellitus, heart diseases, and other complaints. Subjects were divided into different groups with their unique criteria:

1-With the criteria of alcohol consumption: alcoholic (n=13) and nonalcoholic subjects (n=454)2-With the criteria of medication: subjects under medication (n=142) and no medication (n=307)3-With the criteria of diseases: leukemia (n=24), hepatitis (n=10), diabetes mellitus (n=18), and heart diseases (n=27)

Venous blood samples from the subjects were taken into heparinized tubes. CAT activity was measured by the reaction of  $H_2O_2$  decomposing at 240 nm according to the method of Aebi (22). One unit is equal to 1 ?mol of  $H_2O_2$  decomposed/minute. The hemoglobin assay is based on the colorimetric cyanomethemoglobin method and determined by Olympus AU-600 auto analyzer. CAT activity was expressed as k/g Hb. Hematological data that is MCH, HCT, MCHC, MCV, and RBC were studied by Coulter STKS instrument.

Statistical analysis was done by Mann-Whitney U test using SPSS for Windows version 7.5.

#### RESULTS

The results on Table 1 demonstrated that CAT activity of red blood cells from subjects under medication were not significantly different than those of other subjects ?3155 $\pm$ 1039 k/g Hb for subjects under medication and 3051 $\pm$ 956 for other subjects(p>0.05)? CAT activities were 3033 $\pm$ 982 k/g Hb in healthy subjects, 3432 $\pm$ 1126 k/g Hb in

#### Erythrocyte Catalase Activities in Alcohol Consumption, Medications and Some Diseases

	Alcohol		Medication		<b>Diseases</b> Healthy	Leukemia	Heart	Liver	DM
CAT (k/gHb) RBC (10 <sup>6</sup> /µL)	$3644 \pm 985$ 9 $\pm 11$	$3059 \pm 957 \\ 5 \pm 34$	$3155 \pm 1039$ 5 \pm 1	$3051 \pm 957$ $5 \pm 5$	$3033 \pm 982$ $5 \pm 1$	$3432 \pm 1126$ $5 \pm 79$	$2860 \pm 596$ $9 \pm 5$	$3088 \pm 1185$ $5 \pm 1$	$3275 \pm 1126$ $5 \pm 1$
MCV (fL) MCHC	$28\pm 3$ $33\pm 1$	$29\pm 6$ $35\pm 17$	$30{\pm}8$ $35{\pm}8$	$28\pm5$ $35\pm19$	$29\pm 6\ 35\pm 20$	$\begin{array}{c} 29{\pm}10\\ 36{\pm}10\end{array}$	$\begin{array}{c} 32{\pm}13 \\ 35{\pm}9 \end{array}$	$\begin{array}{c} 26{\pm}6\\ 35{\pm}4 \end{array}$	$\begin{array}{c} 28{\pm}1\\ 34{\pm}1 \end{array}$
(g/ aL) HCT (%) MCH (pg)	$\begin{array}{c} 42{\pm}5\\ 84{\pm}5\end{array}$	$\begin{array}{c} 41{\pm}22\\ 87{\pm}54\end{array}$	42±30 83±11	$\begin{array}{c} 40{\pm}17\\ 89{\pm}63 \end{array}$	$\begin{array}{c} 42{\pm}26\\ 89{\pm}64\end{array}$	$39\pm7$ $80\pm7$	$\begin{array}{c} 37{\pm}10\\ 81{\pm}16\end{array}$	$\begin{array}{c} 38{\pm}6\\ 84{\pm}12 \end{array}$	$39{\pm}4$ $83{\pm}3$

Table I. Erythrocyte CAT activity, RBC, MCV, MCHC, HCT, MCH in alcohol consumption, medications and some diseases.

CAT (Catalase), DM (Diabetes Mellitus), RBC (Red Blood Cell), MCV (Mean Corpuscular Volume), MCHC (Mean Corpuscular Hemoglobin Concentration) and HCT (Hematocrit). Data are presented as mean±SD.

leukemia,  $2860\pm596$  k/g Hb in heart diseases,  $3088\pm1185$  k/g Hb in hepatitis, and  $3275\pm1126$  k/g Hb in diabetes mellitus. When the disease groups were compared to the control group, there were no significant differences between them. We found that enzyme activity was affected significantly by only alcohol consumption. CAT activities were  $3059\pm958$  k/g Hb in nonalcoholic subjects and  $3644\pm984$  k/g Hb in alcoholic subjects (p? 0.03).



Figure 1. Erythrocyte catalase activity in our study groups.

## DISCUSSION

The markers were associated with clinical parameters of the disease indicating that reactive oxygen species could play a role in the development of the pathology.<sup>23</sup> In vitro effects of widely used nonsteroidal antiinflammatory drugs (NSAIDs) and paracetamol were studied on oxidative stress-related parameters of human red blood cells (RBC). Erythrocyte CAT activity was increased by Nasalicylate, acemetacin, and tenoxicam at the therapeutic, and by dipyrone at the high concentration.<sup>24</sup>

Some of our findings are inconsistent with data of other authors. This may be partly explained by large differences in the type and size of the populations studied. Differences in the assay conditions (e.g., types of substrates used and precision of measurements) may also affect the result and the degree of significance.

Biochemical changes induced by alcohol in human organism are concentrating especially on the negative influence on the metabolism of liver.25 Alcoholinduced oxidative stress is linked to the metabolism of ethanol. Three metabolic pathways of ethanol have been described in the human body so far.<sup>26</sup> More than 90% of ingested ethanol is metabolized in the body to acetaldehyde and acetate. Ethanol is metabolized in the liver via three distinct enzymatic pathways: alcohol dehydrogenase (ADH), the microsomal ethanol oxidizing system (MEOS) and CAT.<sup>27</sup> Each of these pathways could produce free radicals which affect the antioxidant system. Ethanol per se, hyperlactacidemia and elevated NADH increase xanthine oxidase activity, which results in the production of superoxide. Lipid peroxidation and superoxide production correlate with the amount of cytochrome P450 2E1. MEOS aggravates the oxidative stress directly as well as indirectly by impairing the defense systems. Hydroxyethyl radicals are probably involved in the alkylation of hepatic proteins.<sup>26</sup> The response of the antioxidant defense system in brain subcellular fractions after oral graded doses of ethanol to rat was investigated in a study. Catalase activity was significantly increased in cytosol, synaptosomes and microsomes.<sup>28</sup> In rat thymocytes and cerebellar granule cells, reactive oxygen species (ROS) levels were increased and cell viability was decreased as a result of exposure to ethanol (up to 0.4%).29 Microsomal P450 and peroxisomal fatty acid oxidation activities were studied in liver of rats after long-term ethanol consumption. Ethanol increased peroxisomal beta-oxidation of palmitoyl CoA and CAT activity in liver.<sup>30</sup> It is widely accepted that alcohol metabolism passes through different mechanisms: alcohol dehydrogenase (ADH) activity in stomach epithelial cells, activity of ADH in the liver, MEOS, hepatocyte CAT activity, and

nonoxydizing metabolic pathway (production of fatty acid ethylesters).

We observed a significant increase in erythrocyte catalase activities in alcohol users. In the present study, our observations indicated that increased erythrocyte antioxidant enzyme activities were a possible protective mechanism against oxidative stress induced by alcohol. There are several possibilities here: i) Some of the enzymatic reactions in the glycolysis and pentose phosphate pathway can be destroyed by alcohol or products of alcohol, resulting in a lot of changes in erythrocyte metabolism. ii) Alcohol may enhance molecular oxygen toxicity directly or indirectly, thus, CAT activity may be increased as a compensatory mechanism after these changes. iii) CAT activity may be affected by alcohol but we do not know whether alcohol has inhibitor or activator effects on the CAT enzyme activity. iv) All the antioxidant enzymes in the erythrocyte cytoplasm are in a balance. If one of the factors that affect the antioxidant and oxidant status in the cell is decreased or increase, it results in some minor changes. These changes may contain enzymatic or nonenzymatic systems. In addition, our data suggested that CAT activities were not changed in patients with leukemia, cardiac diseases, hepatitis, and diabetes mellitus according to the control groups.

The data in this study indicate that ethanol ingestion may enhance erytrocyte CAT activity. There was a strong relationship between alcohol drinking and the CAT activity. It was concluded that CAT may be used as an important parameter to determine ethanol induced oxidative stress in erytrocytes.

#### REFERECES

- Carone D, Loverro G, Greco P Lipid peroxidation products and antioxidant enzymes in red blood cells during normal and diabetic pregnancy. Eur J Obstet 1. Gynecol Reprod Biol 1993, 51: 103-109 Agar NS, Sadrzadeh SM, Hallaway PE Erythrocyte catalase. A somatic oxidant
- 2. defense? J Clin Invest 1986, 77: 319-321

- Durak I, Guven T, Birey M Halothane hepatoxicity and hepatic free radical metabolism in guinea pigs. The effects of vitamin E. Can J Anaesth 1996, 43: 741 -3.
- Portoles MT, Catala M, Anton A Hepatic response to the oxidative stress induced by E.coli endotoxin. Glutathione as an index of the acute phase during the endotoxic shock. Mol Cell Biochem 1996, 159: 115-121 4.
- 5. Toborek M, Kopieczna GE, Drozdz M Increased lipid peroxidation and antioxidant activity in methionine-induced hepatitis in rabbits. Nutrition 1996, 12: 534-537
- Bondy SC, Orozco J Effects of ethanol treatment upon sources of reactive 6.
- oxygen species in brain and liver. Alcohol 1994, 29: 375-383 Selden C, Seymour CA, Peters TJ Activities of some free-radical scavenging enzymes and glutathione concentrations in human and rat liver and their 7. relationship to the pathogenesis of tissue damage in iron overload. Clin Sci 1980, 58: 211-219
- Stephens NG, Parsons A, Schofield PM A randomized controlled trial of vitamin 8 CELEVICE 1 AND A CONTRACT OF A
- Oberley LW Free radicals and diabetes. Free Radic Biol Med 1988, 5: 113-124 Wolff S Diabetes mellitus and free radicals. Br Med Bull 1993, 49: 642-652 10.
- L Abbe MR, Trick KD Changes in pancreatic glutathione peroxidase and superoxide dismutase activities in the prediabetic diabete-prone BB rat. Proc Exp 11
- Biol Med 1994, 207: 206-212 12
- Biol Med 1994, 207: 206-212 Roza AM, Pieper GM, Johnston CP Pancreatic antioxidant enzyme activity in normoglycemic diabetes prone BB rats. Pancreas 1995, 10: 53- 58 Salonen JT, Nyyssonen K, Tuomainen TP Increased risk of non-insulin dependent diabetes mellitus at low plasma vitamin E concentrations. A four year 13 study in men. Br Med J 1995, 311: 1124-1127
- Arruda VR, Salles TS, Costa FF Glutathione peroxidase, reduced glutathione, superoxide dismutase and catalase in red cells of patients with hairy cell leukemia 14. Neonlasma 1996 43: 99-102
- 15. Shvalova EP, Antonova TV, Baranovskaia VB The importance of the antioxidant protection systems of the blood in adaptation to the infections process in the infectious process in viral hepatitis B. Ter Arkh 1991, 63: 47-9 Makarenko EV, Kozlovskü IV. The erythrocyte antioxidant system in chronic
- 16. liver disease. Ter Arkh 1989, 61: 115-118
- 17.
- Inver disease. Fer Arkn 1989, 01: 115-118 Inal M, Alatas O, Kanbak G Changes of antioxidant enzyme activities during cardiopulmonary bypass. J Cardiovasc Surg 1999, 40: 373- 376 Zhou J, Du Y, Wang Y The correlation between abusing alcohol and antioxidants, antioxidases. Chung Hua Yu Fang I Hsueh Tsa Chih 1998, 32: 303-18. 305
- 19. Guemouri L, Lecomte E, Herbeth B Blood activities of antioxidant enzymes in
- alcoholics before and after withdrawal. J Stud Alcohol 1993, 54: 626-629 Guemouri L, Arthur Y, Herbeth B Biological variability of superoxide dismutase, glutathione peroxidase, and catalase in blood. Clin Chem 1991, 37: 1932-1937 20.
- Peterson CM, Jones RL, Koenig RJ Reversible hematologic sequelae of diabetes mellitus. Ann Intern Med 1977, 86:425-429 21.
- Aebi H Catalase In: Bergmeyer U, et al Methods of Enzymatic Analysis. New York Academic Press 1974, 673-677 Repetto MG, Reides CG, Evelson P Peripheral markers of oxidative stress in 22. 23
- 24.
- Probable Alzheimer patients. Eur J Clin Invest 1999, 29: 643-649 Orhan H, Sahin G In vitro effects of NSAIDS and paracetamol on oxidative -related parameters of human erythrocytes. Exp Toxicol Pathol 2001, 53: 133-40. Markowski T, Arciuch LP, Zwierz K, Bakush AA Alcoholism: biology. Psychiatr 25.
- Pol 2000, 34: 411-421 26. Zima T, Fialova L, Mestek O, Janebova M, Crkovska J, Malbohan I, Stipek S,
- Mikulikova L, Popov P Oxidative stress, metabolism of ethanol and alcohol-related diseases, J Biomed Sc 2001, 8: 59-70
- Jelski W, Chrostek L, Sznitkowski MMetabolism of ethyl alcohol in the human body. Postepy Hig Med Dosw 1999, 53: 871-883 27.
- Reddy SK, Husain K, Schlorff EC, Scott RB, Somani SM Dose response of ethanol ingestion on antioxidant defense system in rat brain subcellular fractions. 28 Dose Neurotoxicology 1999, 20: 977-987 Huentelman MJ, Peters CM, Ervine WE, Polutnik SM, Johnson P Ethanol has
- 29 differential effects on rat neuron and thymocyte reactive oxygen species levels and cell viability. Comp Biochem Physiol C Pharmacol Toxicol Endocrinol Sep 1999, 124:83-89
- 30 Orellana M. Rodrigo R. Valdes E. Peroxisomal and microsomal fatty acid oxidation in liver of rats after chronic ethanol consumption. Gen Pharmacol 1998. 31: 817-820